PATENT SPECIFICATION

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(54) LIGHT-SENSITIVE COMPOSITIONS

We, HOECHST AKTIENGESELLSCHAFT, a Body Corporate organised according to the laws of the Federal Republic of Germany, of 6230 Frankfurt/Main 80, Federal Republic of Germany, do hereby declare the invention, for which we pray that a patent may be granted to us, and the method by which it is to be performed, to be particularly described in and by the following statement:

This invention relates to light-sensitive compositions which may be supplied either in the form of solid layers or as liquid preparations and which comprise at least one ethylenically unsaturated polymerizable compound and a photoinitiator

system and, optionally, at least one polymeric binder.

Within the term "ethylenically unsaturated polymerizable compounds" there are included low molecular weight polyfunctional monomers capable of addition polymerization and polyunsaturated high molecular weight compounds that can be photocrosslinked.

Substances belonging to the most varied classes of compounds have been proposed as photoinitiators for the photopolymerization of unsaturated compounds. It has been proposed to use as photoinitiators, for example, chalcones, aromatic ketones or diketones, multi-nuclear quinones of the anthraquinone or phenanthrenequinone type, benzanthrone derivatives or aza-benzanthrone derivatives, aromatic nitro compounds, heterocyclic six-membered compounds, such as acridines, phenazines, quinoxalines, quinazolines, pyrylium compounds and thia-pyrylium compounds, heterocyclic five-membered compounds, e.g. benzthiozoles, benzoxazole, or benzimidazoles, and oranic dyestuffs, e.g. eosin, methylene blue or fuchsine.

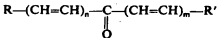
Due to their molecular structure, these photoinitiators very often can be used only for particular layers and substantially lose their effectiveness when used in other polymerizable compositions.

In many cases, it is necessary to use a considerable quantity of the initiator in order to achieve high cross-linking densities, and in these cases the applicability of the initiator is limited by its solubility in the light-sensitive compositions.

Lack of compatibility is frequently observed when the copying composition is

subjected to substantial variations in temperature under unfavorable conditions of storage or transport. Under these conditions, exudation and/or crystallization of the photoinitiator may occur, which result in a loss of light-sensitivity, sometimes considerably impair the adhesion of the light-sensitive layer to a support therefor and substantially detract from the shelf-life of the copying composition.

It has been proposed in German Offenlegungsschrift No. 2060 575, to use unsaturated ketones of the type



40 as photoinitiators in compositions containing monomers with vinyl or vinylidene 40 groups. The light-sensitivity of layers containing only such compounds is not really satisfactory, even if relatrively large quantities of photoinitiator are used nor does their use in combination with special compounds known to be suitable photoconductors from German Patent No. 1 137 625, e.g. appropriately substituted 45 multi-nuclear heterocyclic compounds having groups displaying an electron-donor 45



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effect, meet the demands for high light-sensitivity combined with high cross-linking density made on photoresist materials which must be resistant to corrosive electroplating baths.

1,576,217

It is the object of the present invention to provide a photoinitiator system of good compatibility with the other ingredients of the light-sensitive compositions, which is especially suitable for the photopolymerization of low molecular weight monomeric acrylates and alkylacrylates, and which imparts to the composition good light-sensitivity in combination with a high cross-linking density.

The present invention provides a light-sensitive composition comprising at least one liquid or solid ethylenically unsaturated compound having at least two, preferably terminal, double bonds capable of addition polymerization, other than a compound of the formula II, and a photoinitiator system comprising at least one compound of the general Formula I

wherein

R₀ is O, S, or NR₁,

R₁ denotes hydrogen, a saturated or unsaturated unsubstituted hydrocarbon radical with 1 to 6 carbon atoms, an aminocarbonyl group with from 2 to 11

radical with 1 to 6 carbon atoms, an aminocarbonyl group with from 2 to 11 carbon atoms, an aralkyl group with 7 to 10 carbon atoms, an acyl group with 2 to 18 carbon atoms, an amidomethylene group or a polyalkyleneoxide group of the general formula

$$-(C_0H_{20}-O)_0-C_0H_{20}-OH$$

or the general formula

$$-(C_0H_{20}-O)_0C_0H_{20}-OCH_1$$

wherein a=2 to 4 and n=1 to 10, or a saturated or unsaturated hydrocarbon radical with 1 to 6 carbon atoms which is substituted by hydroxyl, alkoxy, alkoxycarbonyl, acyl, acyloxy, and/or halogen,

R₂ is hydrogen, an alkoxy group, an alkyl group with 1 to 6 carbon atoms, or halogen, and at least one compound of the general Formula II

$$R_3$$
 N
 $(CH=CH)_x - C_1 - R_5$
 R_4
 R_5
 R_5
 R_5

wherein

R₂ is an alkyl group with 1 to 6 carbon atoms, R₄ is an alkyl group with 1 to 6 carbon atoms, and

R₅ is hydrogen, an alkyl group with 1 to 6 carbon atoms, an alkoxy group in which the alkyl group has 1 to 4 carbon atoms, an aryl group with 6 to 12 carbon atoms, or a dialkyl amino styryl group, and wherein is 0 or 1, and, optionally, at least one polymeric binder.

The compounds corresponding to the 6-oxo-anthra(1.9-cd)pyrazole-2(6H) type (Formula I; $R_0=NR_1$) are advantageously substituted in the 2-position (= R_1) by hydrogen, unsubstituted alkyl groups with 1 to 6, preferably 2 to 4 carbon atoms, hydroxyl, alkoxy, alkoxy carbonyl preferably ethoxy carbonyl, acyl or halogen, preferably chlorine or bromine, or by substituted alkyl groups with 1 to 6, preferably 2 to 2 carbon atoms, the numbers 1 to 6 designating the number of carbon atoms arranged in a straight chain in the substituted alkyl group.

Further suitable substituents in the 2-position are: the allyl group, an aralkyl group with 7 to 10 carbon atoms, preferably benzyl or toluyl, or an acyl group with 2 to 18 carbon atoms. Among the acyl groups, acetyl to stearyl, benzoyl, methoxybenzoyl, ethoxy benzoyl, methyl benzoyl, benzsulfonyl, or tosyl are 5 preferred. The substituent R₂ in the six-membered ring of the basic anthracene structure (Formula I; R₀=0, S, NR₂) may be in the 7, 8, 9, or 10-position, preferably in the 10-position, and may be hydrogen, halogen, preferably chlorine or bromine, or 5 alkoxy, preferably methoxy or ethoxy, or alkyl with 1 to 6 carbon atoms, preferably 10 methyl or ethyl. Suitable substituents for R_3 and R_4 in compounds of the type of Formula II are lower unsubstituted alkyl groups with 1 to 6, preferably 1 to 4 carbon atoms, which may be arranged in a straight chain or branched; R_3 and R_4 may be the same or 10 The substituent in R₈ may be hydrogen, a lower alkyl group with 1 to 6, preferably 1 to 4 carbon atoms, an alkoxy group, preferably methoxy, ethoxy, isopropoxy, propoxy or butoxy, an unsubstituted aryl group with 6 to 12 carbon atoms, preferably phenyl, or a substituted aryl group, preferably tolyl, dialkylaminophenyl, or anisyl, or a dialkylamino styryl group. 15 15 20 As indicated above, x may be 0 or 1, but is preferably 0. 20 Compounds corresponding to the Formula I, wherein R_0 stands for N-R₁, may contain, for example, any of the following groups as substituents in the 2-25 -(CH₂)₄-CH₂OH 25 -CH2-CH2-OCH3 -CH2-CH2-OC2H8 $-(CH_2-CH_2-O)_n-CH_2-CH_2OH$ and n = 1-10 $-(CH_2-CH_2-O)_n-CH_2-CH_2OCH_3$ and n = 1-1030 -CH₂-C≡C-CH₂OH 30 -CH₂-CH-CH₂-OCO-CH₃ -CH₂-CH-CH₂-OCO-C₂H₅ 35 -CH2-CH-CH2-OCO-C15H31 35 OH

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$$R_1 = -CH_2 - CH - CH_2 - OCO$$
 wherein $R_3 = H$,
$$OCH_3,$$

$$OC_2H_5,$$

$$-CO - NH$$

$$R_3$$

$$CH_3 \text{ to } C_4H_9$$
 halogen

5 —CO-NH-R₄ wherein
$$R_4 = CH_3$$
, C_2H_5 to C_9H_{19} ,

 $-CH_2-CO-NR_5R_6$ wherein R_5 ; $R_6 = H$, CH_3^1 , C_2H_5 ,

$$-CH_2$$
 wherein $R_7 = OCH_3$, OC_2H_3 , halogen

-CH₂-CH=CH-R₈ and R₈ = CH₃, C₂H₅ -CO-CH=CH.

-CO-CH-CH₂

$$-CH_2-CO-R_9$$
 and $R_9 = CH_{39}$, C_2H_5 to C_6H_{13}

15 and wherein

 $R_2 = H$, OCH_3 , OC_2H_5 or Cl

The following Tables I and II list preferred compounds of the Formulae I and II, respectively:—

TABLE I

No. of the Compound	Substituent R _o	Substituent R,	Substituent R₂
I 1	0		Н
I 2	S	_	· H
I 3	NR ₁	н	Н

TABLE I — Continued.

No. of the Compound	Substituent R _o	Substituent R ₁	Substituent R ₂
I 4	NR ₁	CO—C _e H _e	Н
I 5	NR,	CO—C ₆ H ₄ ·p-OCH ₃	н
I 6	NR,	CO-C ₁₇ H ₃₈	н
. I 7	NR,	CO-C ₇ H ₁₈	Н
I 8	NR,	CO—C ₃ H ₇	Н
I 9	NR,	CO-C ₂ H ₅	Н
I 10	NR,	CO—CH ₃	Н
I 11	NR ₁	SO ₂ C ₆ H ₄ ·p-CH ₃	н
I 12	NR ₁	n-C ₄ H ₉	н
I 13	NR,	i-C ₄ H _e	Н
_ I 14	· NR ₁	Allyl	Н
I 15	NR,	Benzyl	Н
I 16	NR,	CH ₂ —COOC ₂ H ₅	Н
I 17	NR ₁	CH ₂ —CO—C ₆ H ₅	Н
I 18	NR,	CH ₂ CH(OH)CH ₂ CH ₃	н
I 19	NR,	Н	OC ₂ H ₅
I 20	NR,	CO—C ₆ H ₅	OC₂H ₆
I 21	NR ₁	CO-C ₆ H ₄ p-OCH ₃	OC₂H₅
I 22	NR,	CO-C ₁₇ H ₃₅	OC ₂ H ₅
I 23	NR,	CO—C ₇ H ₁₈	OC ₂ H ₅
1 24	NR,	CO—C ₂ H ₅	OC₂H ₈
I 25	NR,	со-сн,	OC ₂ H ₈
I 26	· NR ₁	Benzyl	OC₂H₅
I 27	NR,	CH ₂ —CO—O—C ₂ H ₆	OC₂H₅
I 28	NR,	· CH ₂ —CO—C ₆ H ₅	OC ₂ H ₅
I 29	NR ₁	CH ₂ —CH(OH)—CH ₂ —CH ₃	OC ₂ H ₅
I 30	NR,	Н	Cl
131	NR,	CH ₂ —CH(OH)CH ₂ CH ₃	Cl
I 32	NR,	CH ₂ —CH(OH)—CH ₂ Cl	Cl
I 33	NR ₁	CO—CH ₃	Cl

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TABLE II

No. of the Compound	Subs R ₃	tituents R ₄	Substituent R ₅	x
II 1	СН,	СН,	p-C ₆ H ₄ (CH ₃) ₂	0
II 2	CH ₃	CH ₃	OCH,	0
II 3	CH ₃	CH ₃	OCH ₂ CH ₃	0
II 4	СН,	CH ₃	O-i-C ₃ H ₇	0
II 5	CH ₃	CH ₃	OC ₄ H ₉	0
II 6	CH ₃	СН,	СН,	0
· II 7	СН ₃	СН,	C ₂ H ₅	0
8 11	CH ₃	CH ₃	i-C ₃ H ₇	0
II 9	CH ₃	СН,	$C_{\mathfrak{g}}H_{\mathfrak{g}}$	0
II 1 0	СН ₃	CH ₃	Н	o .
II 11	CH ₃	СН3	$CH=CH-C_6H_4-p-N(CH_3)_2$	1
II 12	CH ₃	CH ₃	$C_{\mathfrak{o}}H_{\mathfrak{o}}$	1
II 13	СН,	CH ₃	p-C ₈ H ₄ OCH ₃	1
II 14	C ₂ H ₅	C ₂ H ₈	C_8H_5	0
II 15	C ₂ H ₅	C ₂ H ₅	$p-C_8H_4N(C_2H_6)_2$	0

Surprisingly, it has been found that, by combining a compound of the Formula I with a compound of the Formula II a light-sensitivity is achieved which in some cases exceeds the sum of the individual effects of the components used by one order of magnitude or more, as expressed, e.g., by the number of fully reproduced steps of a step wedge. That is to say that, in some cases, a synergistic effect is in operation.

Although all compounds of Formula II contain a p-dialkylaminophenyl group, the presence of this group is not alone sufficient to predict a synergistic effect for compounds of this type.

This is demonstrated by the fact, that the compound I 18, for example, does not form a synergistic photoinitiator combination with the following compounds A, B or C. The effectiveness of the compound I 18 is either not increased by these compounds (A, B), or is even reduced (C):

15 A:
$$CH = CH - CH_3$$
 CH_3

B:
$$CH_3$$
 $CH = N$ CH_3

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C:
$$\begin{bmatrix} & & & & & \\ & & & & \\ & & & & \\ & & & & \\ & & & & \\ & & & & \\ & & & & \\ & & & & \\ & & & & \\ & & & & \\ & & & & \\$$

Thus, the marked increase in the photoinitiator effect of the compounds of the Formula I when combined with compounds of Formula II is the more surprising.

The synergistic effect is very pronounced at all mixing ratios, as can be clearly seen from Example 4 below.

The proportions by weight of the compounds of Formula I and Formula II used according to the invention may vary within wide limits and preferably range from 2:98 to 98:2. Proportions between 80:20 and 20:80 are more preferred, and

within these limits proportions from 40:60 to 60:40. Examples of suitable polmerisable compounds are, for example, acrylic and methacrylic acid esters, especially the commercially available ones, for example those of diethylene glycol, triethylene glycol, tetraethylene glycol, polyethylene glycol, trimethylolethane and trimethylolpropane; diglycerol diacrylate, guaiacol glycerol ether diacrylate, neopentyl glycol diacrylate, 2,2-dimethylol butanol-(3)-diacrylate, unsaturated pentaerythritol esters, as described in U.S. Paration and particular pentaerythylol personal glycol and particular pentaerythylol pentaerythylol

3,261,686, reaction products of trimethylol-propane, alkylene oxide and acrylic acids or methacrylic acids in accordance with U.S. Patent No. 3,380,831, and acrylates and methacrylates of polyesters containing hydroxy groups. The latter compounds and other compounds suitable for use in the compositions are described in, for example, U.S. Patents Nos. 2,760,863 and 3,060,023.

The compounds containing urethane groups described in German Offenlegungsschrift No. 2,064,079 and the biuret-group

containing compounds described in German Offenlegungsschrift No. 2,361,041 are also examples of suitable polymerisable compounds.

As will be appreciated the present invention is not limited to the use of specific polymerizable compounds; all that is necessary is that the compound be at least twice ethylenically unsaturated and capable of addition polymerization or be a polyunsaturated, high molecular weight compound that is either capable of being photocrosslinked, or capable of addition polymerization with a low molecular weight compound. The following polyunsaturated compounds, for example, are

suitable for use in light-sensitive compositions:

Polyvinyl cinnamates and prepolymers of unsaturated esters, e.g. the prepolymer of diallyl isophthalate, or the polymeric allyl imides described in German Offenlegungsschrift No. 2,203,732, and the polyvinyl acetals containing extrainear property according to U.S. Patent No. 2,902,710.

The polymerisable compounds preferably contain small amounts (e.g. 50 to ppm) of an inhibitor in order to prevent thermally induced polymerization. If the copying compositions according to the invention are to survive extreme conditions of storage without deterioration, the quantity of inhibitor may be increased to about 1 per cent, calculated on the weight of the polymerisable

compound used. The following compounds may, for example, be used as thermal inhibitors: p-Methoxyphenyl, hydroquinone, alkyl-substituted or aryl-substituted quinones and hydroquinones, tertiary butyl catechol, pyrogallol, copper resinate, naphthyl amine, β -naphthol, copper-(I)-chloride, 2,6-di-tert.-butyl-p-cresol, phenothiazine,

pyridine, nitrobenzene, and dinitrobenzene; p-toluquinone, chloroanil and thiazine dyestuffs, for example Thionine Blue G.

The light-sensitive compositions optionally comprise one or more binders, and normally will do so when the polymerisable compound is a liquid, for example solvent-soluble polyamides, polyvinyl acetates, polymethyl(meth)-acrylates, polyvinyl butyrals, cellulose ethers or cellulose esters, polyalkylene ethers, condensation polymers of glycols with dibasic acids, copolymers of styrene and maleic anhydride which are soluble or at least swellable or softenable in alkaline solutions, copolymers of ethylene and maleic anhydride or of alkyl methacrylates

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	example as described in German Offenle	cribed in German Offenlegungsschrift No. methacrylates, and methacrylic acid, for gungsschrift No. 2,363,806, copolymers of sulfonyl)-carbamic acid-(\beta-methacryloxy)-	
5	further malezate resins, terpenphenol re Aqueous alkaline developers are	erman Offenlegungsschrift No. 2,027,466, esins, and others. preferably used for development and	. 5
10	and maleic anhydride and copolymers of terpolymers of styrene, alkyl methacrylat the copolymers described in German O	are either alkali-soluble or softenable in such binders are copolymers of styrene alkyl methacrylate and methacrylic acid, e and methacrylic acid, maleic resins, and ffenlegungsschrift No. 2,205,046.	10
15	absorbants, dyestuffs, pigments, color sensitometric regulators, for example, minvention.	composers, hydrogen donors, oxygen couplers, UV-absorbing substances and ay be included in the compositions of the	15
20	taken that the substances added do not a light required for the initiating processensitivity of the copying composition	bsorb an undue proportion of the actinic s and thus reduce the practical light-	20
25	aliphatic polyols. If good storability at hig water-insoluble plasticizers are preferable	th atmospheric moisture is to be ensured, y used. added in cases where the light-sensitive	25
30	been found to be advantageous in such cisilanes, nitrogen-containing heterocyclic in U.S. Patents No. 3,645,722, No. 3,6 mercaptans, for example those describe	The following adhesion promoters have ases:— monomeric or polymeric organic compounds, for example those described 22,234 and No. 3,827,908, heterocyclic and in German Offenlegungsscheit No.	30
35	Offenlegungsschrift No. 2,448,821, and me those described in German Offenlegungs	for example those described in German ercapto alkanoic acid esters, for example schrift No. 2,448,850. Example, substances containing aliphatic pounds. In some cases, the hinder or the	35
40	contains a labile hydrogen atom. Further, the compositions of the investigation of the invest	ention may comprise dyestuffs and/or	40
45	The light-sensitive copying composition U.V. absorbers serving as anti-halation filted described, for example, in German Offen	legungsschrift No. 2,243,182. omponents of the light-sensitive copying es heing by weight and being calculated	45
50	binder:	15—99 per cent	50
	monomer:	1—75 per cent	
	photoinitiator mixture:	0.1—10 per cent	
	hydrogen donor:	0.5—10 per cent	
	plasticizer:	0.0—15 per cent	
55	adhesion promoter:	0.0—15 per cent	55
•	dyestuff or pigment:	0.0—30 per cent.	

	For commercial purposes, the light-sensitive copying compositions may be supplied in the form of solutions or dispersions, i.e. as so-called photoresist	
	lacquers, which the customer himself applies to a particular support — this being the usual procedure, for example, for chemical milling, for the preparation of	
5	printed circuits or stencils, or for making printing plates and screen printing stencils — followed by drying, imagewise exposure and development. In this case,	. 5
	the ingredients of the light-sensitive copying composition are dissolved in a suitable solvent. Suitable solvents are, for example, alcohols, ketones, esters, ethers, amides	
10	and hydrocarbons. The partial ethers of glycols or of keto-alcohols have proved to be particularly favorable solvents but the choice of solvent is substantially	10
•	determined by the binder, if any, present in the composition. Alternatively, the light-sensitive copying compositions may be marketed in the	
	form of a solid photopolymerizable layer disposed on a support and may be used as such for the preparation of, for example, printing forms, relief images, etch resists.	
15	stencils, matrices, screen printing forms or single copies. A particularly important application is as storable pre-sensitized printing plates processable into	15
•	planographic, relief or intaglio printing formes. The supports are coated with the compositions from appropriate organic	
20	solvents or solvent mixtures, using, for example, casting, spraying or immersion processes. Suitable supports comprise, for example, magnesium, zinc, copper	20
	mechanically, chemically or electrochemically roughened aluminium, anodized aluminium, steel, polyester or acetate films, polyamide, e.g., "Perlon" gauze, the	
25	surfaces of which may be subjected to a pre-treatment, where necessary. The support may serve either as the final, permanent, support or as an intermediate	
25	support from which the light-sensitive copying composition is transferred, by lamination, onto the workpiece to be processed. If thick photopolymer layers are	25
	to be produced, i.e. of a thickness of several tenths of a millimetre, the copying composition may be kneaded without dissolution, for example in a three-roll mill, and then hydraulically record onto the system for example in a three-roll mill,	
30	and then hydraulically pressed onto the support, for example for one minute at a pressure of 30,000 to 50,000 kp and a temperature of 90°C. If crosslinking is effected solely by polymerization of the ethylenically	30
	unsaturated compound(s), it is normally of advantage to protect the lightsensitive copying composition, during exposure, from atmospheric oxygen, because oxygen	•
35	tends to intercept the radicals forming within the layer and to deactivate them. Such an exclusion of oxygen may be achieved by providing the light-sensitive layer	35
	with an oxygen-impermeable barrier layer, as is described, for example, in German Offenlegungsschriften Nos. 1,572,153 and 2,036,585.	
40	If the light-sensitive copying composition contains a high molecular weight compound which is capable of a photochemical crosslinking reaction, so that	
40	crosslinking is not exclusively or predominantly caused by, for example, low molecular weight acrylates or alkyl acrylates, no such barrier layer is necessary,	40
	because in this case photo-crosslinking results in the desired differentiation between exposed and unexposed areas even in the presence of oxygen.	
45	Material prepared with the light-sensitive copying compositions according to the invention may be used, on the one hand, for the preparation of images on suitable support or receiving sheets, and, on the other hand, for the preparation of	45
	relief images which may be used, for example, as printing forms, screens, and photoresists. In addition, it is possible for the light-sensitive copying compositions	
50	of the present invention to be used for the preparation of UV-hardening lacquers which may be used for surface protection, or for the preparation of UV-hardening	50
	printing inks which do not dry physically and do not cross-link chemically, with formation of cross-connections by oxygen action. Drying is by a photochemical	
55	process and thus is particularly fast and ecologically safe. Printing forms, screens, and etch resists may be prepared from appropriate	
33	copying materials according to the invention in the usual manner, e.g. after exposure to actinic light through a suitable original the non-image areas are	55
	removed by treatment with suitable solvents, e.g. aqueous alkaline solutions, in which those areas are soluble but in which the image areas, by virtue of their being	
60	hardened by the action of the light, are insoluble. Alternatively, development may be effected by other methods, utilizing other physical differences between the hardened image and the unit to the physical differences between the hardened image.	60
	physical differences between the hardened image areas and the unhardened non- image areas, for example differences in their melting points, their stickiness, their adhesiveness or their optical transparency.	
	The invention is illustrated by some of the following Examples. Other	

10	1,576,217	10
	Examples are included for comparison purposes. Unless otherwise stated, all quantities are by weight. The relation between parts by weight and parts by volume is the same as that between grams and milliliters.	
5	Example 1. 140.0 p.b.w. of a copolymer of 85.8 p.b.w. of methylmethacrylate and 12.5 p.b.w. of methacrylic acid with an average molecular weight of 35,000 and an acid number of 86,	5
10	140.0 p.b.w. of pentaerythritol-triacrylate, and 1.5 p.b.w. of tri-[4-(methyl-phenylamino)-phenyl]-methyl acetate are dissolved in 1400.0 p.b.w. of ethylene glycol monoethyl ether.	10
15	A series of solutions of the above composition are prepared, and in each case either 7.0 p.b.w. of one initiator, or a mixture of 3.5 p.b.w. of an initiator according to Formula I and 3.5 p.b.w. of an initiator according to Formula II are added. The solutions are filtered and then whirler-coated, at 100 revolutions per minute, onto an electrochemically roughened aluminum foil and dried. After drying, the sample plates are so coated with an aqueous solution containing	15
20	2.0 p.b.w. of carboxy methyl cellulose, 1.0 p.b.w. of cane sugar, 1.0 p.b.w. of saponin, and 0.12 p.b.w. of sorbic acid, in 267.0 p.b.w. of water,	20
25	that the dry layer weighs about 0.6 g per square meter. Then each sample is exposed for two minutes under a 21-step half-tone wedge with a density range of 0.05—3.05 and density increments of 0.15 (Kodak*: Photographic Step Tablet No. 2). The light source used is an 8.000 W "Xenokon" and are large (distance 7.2).	25
30	For removal of the non-image areas, the plates are wiped over for 30 seconds with a developer having a pH of 11.9 and being composed of: 1.5 p.b.w. of sodium metasilicate nonahydrate, 0.3 p.b.w. of polyethylene glycol 6000, 0.06 p.b.w. of levulning acid, and 0.03 p.b.w. of stronting acid, and	30
35	0.03 p.b.w. of strontium hydroxide octahydrate in 100.0 p.b.w. of water	35
40	and then rinsed down with water. If the copying layers are processed in this manner, the fully reproduced steps of the Kodak step wedge are a measure for the starter activity of the initiators or initiator mixtures tested, i.e. the higher the number of steps reproduced, the higher the practical light-sensitivity. The following Table III lists the number of fully reproduced steps of the step wedge for each of the initiator combinations tested, adjacent partially colored transition steps being disregarded. The light-sensitivities of two adjacent steps of the step wedge differ by a factor of A. Sens O entitle the step wedge differ by a factor of A. Sens O entitle the step wedge differ by a factor of A. Sens O entitle the step wedge differ by a factor of A. Sens O entitle the step wedge differ by a factor of A. Sens O entitle the step wedge differ by a factor of A. Sens O entitle the step wedge differ by a factor of A. Sens O entitle the step wedge differ by a factor of A. Sens O entitle the step wedge differ by a factor of A. Sens O entitle the step wedge differ by a factor of A. Sens O entitle the step wedge differ by a factor of A. Sens O entitle the step wedge differ by a factor of A. Sens O entitle the step wedge differ by a factor of A. Sens O entitle the step wedge differ by a factor of A. Sens O entitle the step wedge differ by a factor of A. Sens O entitle the step wedge differ by a factor of A. Sens O entitle the step wedge differ by a factor of the step wedg	40
45	the step wedge differ by a factor of $\sqrt{2}$. Step 0 of the step wedge corresponds to an optical density of 0.05 (inherent absorption of the film material).	45

^{* &}quot;Kodak is a trade mark

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	Initiators	ABLE III	
Type I	Type II	Test No.	Number of Steps
1 9		1	1
I 14		2	1
I 15		3	· 3
	II 1	4	1
	II 3	. 5	io
	II 6	6	0
	II 9	7	0
	II 15	8	1
I 9	II 1	9	7
I 9	II 3	. 10	8
I 9	II 6	11	9
I 9	II 9	12	6
I 9	II 15	. 13	3
I 14	II 1	14	6
I 14	II 3	15	10
I 14	II 6	16	10
I 14	II 9	17	7
I 14	II 15	18	. 5
I 15	II 1	19	8
I 15	II 3	20	8
I 15	II 6	21	8
I 15	II 9	22	9
I 15	II 15	23	4

Example 2.

This example shows the influence of different compounds of Formula II on a specific compound of Formula I (Compound No. I 18).

140.0 p.b.w. of a copolymer of 85 p.b.w. of methyl methacrylate and 15 p.b.w. of methacrylic acid with an average molecular weight of 43,000 and an acid number of 86,

140.0 p.b.w. of trimethylol ethane triacrylate, and
1.5 p.b.w. of a blue azo dyestuff, obtained by coupling 2,4-dinitro-6-chlorobenzene-diazonium salt with 2-methoxy-5-acetylamino-N-cyano-ethyl-N-10 hydroxy-ethyl aniline, are dissolved in

1400.0 p.b.w. of ethylene glycol monoethyl ether.

A series of such solutions are prepared and to each of them either 5.0 p.b.w. of

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one initiator, or 2.5 p.b.w. of the initiator corresponding to Formula I 18 and 2.5 p.b.w. of one of the initiators corresponding to Formula II are added.

The solutions are filtered and whirler-coated, at 100 revolutions per minute, onto an electrochemically roughened and anodized aluminum foil.

The dried samples are coated with a solution of 3.5 p.b.w. of polyvinyl alcohol and 1.0 p.b.w. of sodium lauryl-ether-sulfate in

96.5 p.b.w. of water

which has a viscosity of about 16 cp and are then dried. Further processing is as described in Example 1. Table IV indicates the relative light-sensitivities of the samples obtained.

TABLE IV

	Initiators	LE IV			
Type I	Type II	Test No.	Number of Steps		
I 18		24	1		
	II 14	25	0		
	II 11	26	0		
	II 2	27	no image		
	II 15	28	0		
	II 12	29	0		
	II 13	30	0		
	II 6	31	0		
I 18	II 14	32	7.		
I 18	П 11	33	4		
I 18	II 2	.34	12		
I 18	II 15	.35	8		
I 18	II 12	. 36	5		
I 18	II 13	37	6		
I 18	II 6	38	11		

Example 3.

This example serves to show the influence of one initiator compound of Formula II upon different initiator compounds of Formula I.

The tests were carried out analogously to Example 2. Several compounds were used as initiators corresponding to Formula I, and the Initiator of Formula II was the compound numbered II 1.

The results are listed in the following Table V.

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	Initiators	•	TABLE V	
Type I	imuators	Type II	Test No.	Number of Steps
I 1			39	1
I 9			40	5
I 17			41	4"
I 18			42	1
I 28			43	0
		II 1	44	1
1 1	·	II 1	45	10
I 9		II 1	46	12
· 117		II 1	47	12
I 18		Ii 1	48	_11
I 28		II 1	49	8

Example 4.

This example shows how wide the limits in which the mixing ratio of the two components of the initiator system may be varied without losing its synergistic

140.0 p.b.w. of a copolymer of 85 p.b.w. of methyl methacrylate and 15 p.b.w. of methacrylic acid with an approx. average molecular weight of 43,000, 140.0 p.b.w. of trimethylol ethane triacrylate, and 1.5 p.b.w. of the blue dyestuff used in Example 2

are dissolved in

1400.0 p.b.w. of ethylene glycol monoethyl ether.

A series of such solutions are mixed in each case either with 5.0 parts by weight of the initiator numbered I 1 or the initiator numbered II 1, or with 5.0 parts by weight of a mixture of these two initiators.

The tests are carried out exactly as the tests described in Examples 2 and 3. The results are listed in the following Table VI.

TABLE VI

Initi Type I	ators Type II	Test No.	Number of Steps
5.0 p.b.w.		39	1
4.0 p.b.w.	1.0 p.b.w.	50	. 8
3.0 p.b.w.	2.0 p.b.w.	51	9
2.5 p.b.w.	2.5 p.b.w.	45	10
2.0 p.b.w.	3.0 p.b.w.	52	11
1.0 p.b.w.	4.0 p.b.w.	53	10
0.5 p.b.w.	4.5 p.b.w.	54	9
0.25 p.b.w.	4.75 p.b.w.	55	8
	5.0 p.b.w.	44	1

	1,370,217	14
5	Example 5. In this example, compounds according to Formula I carrying different substituents in the 10-position are compared. 140.0 p.b.w. of a copolymer of 87.5 p.b.w. of methacrylate and 12.5 p.b.w.	
	140.0 p.b.w. of trimethylol ethane triacrylate, and 1.5 p.b.w. of the blue dyestuff described in Ex. 2 are dissolved in	5
10	1400.0 p.b.w. of ethyleneglycol monoethyl ether. A series of such solutions are mixed either with 5.0 p.b.w. of one initiator, or with 2.5 p.b.w. each of two initiators. The samples are evaluated as in Example 1 and the results are listed in the following Table VII.	10

TABLE VII

Initi Type I	iators Type II	Test No.	Number of Steps
I 18		56	1
I 18	II 1	57	9
I 29		58	no image
I 29	II 1	59	4
I 31		60	1
I 31	II 1	61	6
	II 1	62	1

15	Example 6.	
	This example serves to illustrate that even in light appointment	15
	The state of the s	
	the synergistic effect of the initiator systems according to the present invention is fully effective.	
20	A series of solutions are prepared comprising	
	300.0 p.b.w. of diallyl isophthalate prepolymer and	20
	90.0 p.b.w. of pentaerythritol triacrylate in	
	2900.0 p.b.w. of 2-methyl-2-methoxynentanone-4	
25	Each of these solutions is mixed either with 16.0 p.b.w. of a single initiator, or with 8.0 p.b.w. each of the two initiator.	
	with 8.0 p.b.w. each of the two initiators of Formula I and Formula II to be used in mixture, and the solutions are then agitated until the solid substances have	25
	completely dissolved. The solillions are filtered and while an account of the	
	TO TO THE POPULATION OF THE PO	
30	are minum tons are unicular to minumes at any and are then evaced in a second	
-	maine under a 21-step han-tone step wedge (k odak, photographic care Tell 1 2)	30
	-// wing an ovve w Action Soul are lamb from a distance of 72 am	50
	After exposure, the samples are bathed for 60 seconds in 1.1.1-trichloroethane and then sprayed with clean solvent. The plates are then treated for 45 seconds, with the aid of a cotton and with the sid of a cotton and w	
35	with the aid of a colloil day, with the etching collition described in Common one	
33	iogungssemmit 140. 1.740.200 and comprising.	25
	80.0 p.b.v of gum arabic (14°Be)	35
	12.0 p.b.v. of phosphoric acid (85%) 0.2 p.b.v. hydrofluoric acid (50%),	
	0.5 p.b.v. of H_2O_2 (30%), and	
40	7.3 p.b.v. of water,	
	The image areas are then inked up with greasy ink. The relative to the	40
	oversity ness are compared in the same manner as described in Evample 1	
	The results of the evaluation are listed in the following Table VIII	

TABLE VIII Initiators						
Туре І	Type II	Test No.	Number of Steps			
	II 1 (8.0 p.b.w.)	63	8			
	II 1 (16.0 p.b.w.)	64	8			
I 2		65	4			
I 2	II i	66	13			
I 4		67	7			
I ,4	II 1	68	14			
I 6		69	6			
I 6	II I	70	15			
1 7		71	5			
I 7	II 1	72	13			
· I 8		73	8			
I 8	II 1	74	13			
I 9		75	8			
I 9	II 1	76	15			
I 12	1	77	3			
I 12	II 1	78	14			
I 13	•	. 79	3			
I 13	II 1 ·	80	13			
I 15		81	8			
I 15	II 1	82	15			
I 16		83	7			
I 16	п	84	16			
I 17		85	7			
I 17	. II 1	86	15			
I 24		87	3			
I 24	II 1	88	13			
I 29		89	3			
I 29	II 1	90	13			
1 30		91	i			
	II 1	92	14			
I 33		93	6			
-1 33	II 1	94	16			

and dried. The coated fabric is image-wise exposed for 4 minutes under a positive

original, using an 8 kW xenon lamp at a distance of 72 cm, and the resulting image

is developed with acetone. A useful silk screen is thus obtained.

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	-,0.0,017	17
	Example 10.	
	A coating solution is prepared from	
	400.0 p.b.w. of styrene/maleic anhydride copolymer, 200.0 p.b.w. of an α-methylstyrene/vinyltoluene copolymer,	
5	25.0 p.b.w. of a polyvinyl butryal with an average molecular weight of 30,000 to	_
	33,000,	5
	500.0 p.b.w. of trimethylol propanetriacrylate,	
	1.0 p.b.w. of Compound I 1, and	
10	4.0 p.b.w. of Compound II 1, in	
	2000.0 p.b.w. of methyl ethyl ketone and 200.0 p.b.w. of ethyleneglycol monoethyl ether	10
	and filtered.	
	A 35 μ m thick copper foil laminated to a base of phenolic resin paper is coated with this solution. After draine the leave to a base of phenolic resin paper is coated	
	with this solution. After diving, the laver is covered with a 1 to 2 um thick not make in the	
15	arconor layer,	15
	The coated copper plate is exposed for 2 minutes under a negative original	13
	showing a circuit didgitalli, using an a kw yenon lamn. Then it is developed with the	
	aqueous aixainie developer used in Example I and etched at 5000 with an aqueous	
20		
	The copper is etched away in the bared areas and a printed circuit of the type	20
	used in the electrical industry is obtained.	
	Example 11.	
	700.0 p.b.w. of a terpolymer of 25 p.b.w. of methacrylic acid, 62.5 p.b.w. of n-	
	incommendation viale, and 12.3 p.p.w. of methylmethacrylate as a hinder	
25	500.0 p.o.w. of the monomer described in his 1 of German Offenlesungeschaft	25
	100 2007,077 HEACHOIL DECORET OF 22 Astrimethylhogomethylans	23
	ansocyanate and b-nydroxy-einvimernacrylare)	
	13.0 p.b.w. of trietnyleneglycol dimethacrylate	
30	2.0 p.b.w. of the blue dyestuff used in Ex. 2,	
	5.0 p.b.w. of Compound I 28, and 5.0 p.b.w. of Compound II 6	30
	are dissolved in	
•	2500.0 p.b.w. of methyl ethyl ketone and	
	200.0 p.b.w. of ethyl alcohol.	
35	The solution is filtered and coated onto a polyethylene terephtholate annual	35
	in such a manner that the resulting ary laver weache 17 a/m2	33
	Dy means of a commercially available laminating apparatus, the control classic	
	adminuted onto the copper surface of a composite material consisting of larges of	
40	copper and naturpaper.	
	After a two minutes' exposure of the laminate (xenon lamp, 8 kW) under a	40
	positive original showing a circuit diagram, the protective film is pulled off and the image of the circuit diagram is developed with a 0.8 per cent strength sodium	
	carbonate solution.	
45	The copper circuit paths are reinforced by electrodensition and the	
45	overed, also by electrodeposition, with a lead-tin laver	45
	Inc resist layer is removed by immersion in acetone. Finally, the board	73
	is stoned away in a 1 CC12 SULLIUII. HILLS COMPLETING the photochemical meaduration	
50	In our copending Application No. 54130/76 (Serial No. 1,576,218) there is	
		50
	unsaturated polymerizable compound, a photoinitiator of the formula I, as given above, and optionally a binder.	
	assis, and optionally a binder.	
	WHAT WE CLAIM IS:—	
55	I. A light-sensitive composition composition as least the sensitive composition composition as least the sensitive composition as least the sensitive composition composition as least the sensitive composition composition as least the sensitive composition composition composition as least the sensitive composition com	
		EE
		55
	system comprising at least one compound of the general Formula 1	
		•

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wherein

R_o is O, S or NR,

R, denotes hydrogen, a saturated or unsaturated unsubstituted hydrocarbon radical with 1 to 6 carbon atoms, an aralkyl group with 7 to 10 carbon atoms, an acyl group with 2 to 18 carbon atoms, an amidomethylene group, an aminocarbonyl group with from 2 to 11 carbon atoms, a polyalkyleneoxide group of the general formula

 $-(C_aH_{2a}-O)_n-C_aH_{2a}-OH$

10 or the general formula

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$$-(C_aH_{2a}-O)_n-C_aH_{2a}-OCH_3$$

wherein a=2 to 4 and n=1 to 10, or a saturated or unsaturated hydrocarbon radical with 1 to 6 carbon atoms which is substituted by hydroxyl, alkoxy, alkoxycarbonyl, acyl, acyloxy, and/or halogen,

R₂ is hydrogen, an alkoxy group, an alkyl group with 1 to 6 carbon atoms, or halogen, and at least one compound of the general Formula II

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$$\begin{array}{c|c}
R_3 & & \\
R_4 & & \\
\end{array}$$

$$N \longrightarrow \begin{pmatrix} CH=CH \\ \times & -C \\ 0 & \\
\end{array}$$

$$CH=CH \times \begin{pmatrix} CH=CH \\ \times & C \\ 0 & \\
\end{array}$$

wherein

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R₃ is an alkyl group with 1 to 6 carbon atoms,
R₄ is an alkyl group with 1 to 6 carbon atoms, and
R₅ is hydrogen, an alkyl group with 1 to 6 carbon atoms, an alkoxy group in which the alkyl group has 1 to 4 carbon atoms, an aryl group with 6 to 12 carbon atoms, or a dialkyl amino styryl group, and wherein

is 0 or 1. and optionally, at least one polymeric binder.

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2. A composition as claimed in claim 1, wherein the double bonds in at least one ethylenically unsaturated compound are terminal double bonds.

3. A composition as claimed in claim 1 or claim 2 wherein the initiator comprises at least one of the specific compounds of the general formula I mentioned herein.

4. A composition as claimed in any one of claims 1 to 3 wherein the initiator comprises at least one of the specific compounds of the general formula II

mentioned herein.

5. A composition as claimed in any one of claims 1 to 4 wherein the compounds of the general formula I and general formula II are present in a weight ratio of from 2:98 to 98:2.

6. A composition as claimed in any one of claims 1 to 5 comprising, as a polymerisable compound, an acrylic or methacrylic acid ester.

7. A composition as claimed in claim 6 comprising at least one of the acrylic or

methacrylic esters specified herein. 8. A composition as claimed in any one of claims 1 to 7 comprising, as a

polymerisable compound, a high molecular weight, polyunsaturated, polymerisable

9. A composition as claimed in any one of claims 1 to 8 which contains a polymeric binder which is softenable or soluble at room temperature in an aqueous alkaline solution.

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10. A composition as claimed in claim 9 wherein the binder comprises a styrene/maleic anhydride copolymer, an alkyl methacrylate/methacrylic acid	
11. A composition as claimed in any one of claims 1 to 10 comprising a thermal	_
12. A composition as claimed in claim 1 substantially as described in any one of tests 9 to 23, 32 to 38, 45 to 49, 50 to 55, 57, 59, 61, 66, 68, 70, 72, 74, 76, 78, 80, 82,	5 ,
84, 86, 88, 90, 92 and 94 herein or in any one of Examples 7 to 11 herein. 13. A composition as claimed in any one of claims 1 to 12 in the form of a	10
14. A composition as claimed in any one of claims 1 to 13 in the form of a solid	10
15. A composition as claimed in claim 14 wherein the carrier comprises	
16. A composition as claimed in claim 14 or claim 15 wherein the surface of the layer remote from the carrier has thereon an oxygen-impermeable protective layer. 17. A composition as claimed in claim 14 substantially as described in any one	15
18. A method of making an article comprising at least the steps of image-wise exposing to actinic light a composition according to claim 16 and then processing	20
unexposed portions of the layer.	
treated with a liquid developer to dissolve or soften the unexposed portions of the layer.	25
20. An article whenever made by the method of claim 18 or claim 19. 21. An article as claimed in claim 20 which is a printing forme. 22. An article as claimed in claim 20 which is a printed circuit board.	
	styrene/maleic anhydride copolymer, an alkyl methacrylate/methacrylic acid copolymer or a maleate resin. 11. A composition as claimed in any one of claims 1 to 10 comprising a thermal polymerisation inhibitor. 12. A composition as claimed in claim 1 substantially as described in any one of tests 9 to 23, 32 to 38, 45 to 49, 50 to 55, 57, 59, 61, 66, 68, 70, 72, 74, 76, 78, 80, 82, 84, 86, 88, 90, 92 and 94 herein or in any one of Examples 7 to 11 herein. 13. A composition as claimed in any one of claims 1 to 12 in the form of a solution or dispersion. 14. A composition as claimed in any one of claims 1 to 13 in the form of a solid layer on a carrier. 15. A composition as claimed in claim 14 wherein the carrier comprises aluminium, magnesium, zinc, copper, steel or a plastics material. 16. A composition as claimed in claim 14 or claim 15 wherein the surface of the layer remote from the carrier has thereon an oxygen-impermeable protective layer. 17. A composition as claimed in claim 14 substantially as described in any one of the Examples herein. 18. A method of making an article comprising at least the steps of image-wise exposing to actinic light a composition according to claim 16 and then processing the exposed composition in order to remove either the exposed portions or the unexposed portions of the layer. 19. A method as claimed in claim 18 wherein the exposed composition is treated with a liquid developer to dissolve or soften the unexposed portions of the layer. 20. An article whenever made by the method of claim 18 or claim 19. 21. An article as claimed in claim 20 which is a printing forme.

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